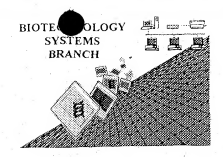
RAW SEQUENCE LISTING ERROR REPORT



The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number: 09/673, 840

Source: PCT 09

Date Processed by STIC: 2-15-01

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT.

2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT,

FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 c-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 c-mail help: patin3help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE CHECKER VERSION 3.0 PROGRAM, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW:

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings generated for the original version of 37 CFR §§1.821 – 1.825 effective October 1, 1990 (old rules) and the revised version (new rules) effective July 1, 1998 as well as World Intellectual Property Organization (WIPO) Standard ST 25

Checker Version 3.0 replaces the previous DOS-based version of Checker, and is Y2K-compliant. Checker allows public users to check sequence listings in Computer Readable form (CRF) before submitting them to the United States Patent and Trademark Office (USPTO). Use of Checker prior to filing the sequence listing is expected to result in fewer errored sequence listings, thus saving time and money

Checker Version 3.0 can be down loaded from the USPTO website at the following address: http://www.uspto.gov/web/offices/pac/checker

SERIAL NUMBER:

ERROR DETECTED SUGGESTED CORRECTION

1	N: NEW RULES CASES: _ Wrapped Nucleics	PLEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE The number/text at the end of each line "wrapped" down to the next line. This may occur if your file was retrieved in a word processor after creating it.
	A second	Please adjust your right margin to 3, as this will prevent "wrapping".
2	_ Wrapped Aminos	The amino acid number/text at the end of each line "wrapped" down to the next line. This may occur if your file was retrieved in a word processor after creating it. Please adjust your right margin to .3, as this will prevent "wrapping".
3	_ Incorrect Line Length	The rules require that a line not exceed 72 characters in length. This includes spaces.
4	_ Misaligned Amino Acid Numbering	The numbering under each 5th amino acid is misaligned. This may be caused by the use of tabs between the numbering. It is recommended to delete any tabs and use spacing between the numbers.
5	_ Non-ASCII	This file was not saved in ASCII (DOS) text, as required by the Sequence Rules. Please ensure your subsequent submission is saved in ASCII text so that it can be processed.
6	Variable Length	Sequence(s) contain n's or Xaa's which represented more than one residue. As per the rules, each n or Xaa can only represent a single residue. Please present the maximum number of each residue having variable length and indicate in the (ix) feature section that some may be missing.
7	Patentin ver. 2.0 "bug"	A "bug" in Patentin version 2.0 has caused the <220>-<223> section to be missing from amino acid sequence(s) Normally, Patentin would automatically generate this section from the previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section to the subsequent amino acid sequence. This applies primarily to the mandatory <220>-<223> sections for Artificial or Unknown sequences.
8	Skipped Sequences (OLD RULES)	Sequence(s) missing. If intentional, please use the following format for each skipped sequence: (2) INFORMATION FOR SEQ ID NO:X: (i) SEQUENCE CHARACTERISTICS:(Do not insert any headings under "SEQUENCE CHARACTERISTICS") (xi) SEQUENCE DESCRIPTION:SEQ ID NO:X: This sequence is intentionally skipped Please also adjust the "(iii) NUMBER OF SEQUENCES:" response to include the skipped sequence(s).
9	Skipped Sequences (NEW RULES)	Sequence(s) missing. If intentional, please use the following format for each skipped sequence. <210> sequence id number <400> sequence id number 000
10	Use of n's or Xaa's (NEW RULES)	Use of n's and/or Xaa's have been detected in the Sequence Listing. Use of <220> to <223> is MANDATORY if n's or Xaa's are present. In <220> to <223> section, please explain location of n or Xaa, and which residue n or Xaa represents.
11	Use of <213>Organism (NEW RULES)	Sequence(s) are missing this mandatory field or its response.
12	Use of <220>Feature (NEW RULES)	Sequence(s) are missing the <220>Feature and associated headings. Use of <220> to <223> is MANDATORY if <213>ORGANISM is "Artificial" or "Unknown" Please explain source of genetic material in <220> to <223> section. (See "Federal Register," 6/01/98, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of new Rules)
13	Patentin ver. 2.0 "bug"	Please do not use "Copy to Disk" function of Patentin version 2.0. This causes a corrupted

Instead, please use "File Manager" or any other means to copy file to floppy disk.

Does Not Comply Corrected Diskette Needed

PCT09

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/673,840

DATE: 02/15/2001

TIME: 14:19:53

Input Set : A:\ES.txt

Output Set: N:\CRF3\02152001\1673840.raw

- 5 <110> APPLICANT: metaGen Gesellschaft f?r Genomforschung mbH (Assignee)
- 7 <120> TITLE OF INVENTION: Human Nucleic Acid Sequences from Normal Bladder Tisssue
- 9 <130> FILE REFERENCE: 51587AWOM1XX24-P
- C--> 11 <140> CURRENT APPLICATION NUMBER: US/09/673,840
- C--> 12 <141> CURRENT FILING DATE: 2000-10-23
 - 14 <160> NUMBER OF SEQ ID NOS: 431

ERRORED SEQUENCES									Invalid amino acid designator											
			<210> <211>																vences	
			<212>							Co	ntai	ning	Xa	a's '	" or	n'5	" n	1057	have	
		11009	<213>	ORGAN	IISM:	homo	sapie	ens										-		
		11011	<400>	SEQUE	INCE:	416	_			< 2	207	To 4	22	5/	yea:	701G	٠	70	explain	
	E>	11013	Tyr	Thr	Met	(XXX)	lle	Ile	Tyr	Phe	Thr	Arg(Xxx) Ile	Leu	Tyr	(xxx)	Gln	what they	
		11014	1				5		-			10					15		',	
	E>	11016	Gly	Gly	Ile	Leu	Lys	Tyr	Asn	Thr	Pro	Gly	(xxx)	Ser	Phe	Leu	Leu	$\mathbf{T}\mathbf{y}\mathbf{r}$	represent.	
		11017				20					25					30			<i>*</i> ,	
	E>	11019	Ile	Met	Ile	Val	Ser	Phe	His	Ile	Ser	Trp	(xxx)	Leu	(xxx)	(XXX)	Gly	Lys	(They may	
		11020			35					40			-		45				only represe	
	E>	11022	${\tt Gly}$		(XXX	🕽 Lys	Ser	Ile	Phe	Ile	${ t ryr}$	Ile	Lys	Thr	Lys	(xxx)	(xxx)	Gln	only represe	200
		11023		50					55					60					a single	
	E ~ - >	11025	182.00	Arg	Leu	(XXX)	Pro	Pro	${ t Lys}$	Cys	Leu	Val		Leu	Glu	Asn	Asn	Met	residue)	
		11026	65		- www.	and the same of th		7,0					75					80	1 CSIONE J	
		11028	Asn	Gluc	(XXX)	(XXX)	_	Met	Asn	Gln	Ile	Thr	Trp(Xxx) Thr	His	Arg	Arg		
		11029				-	85	_	_	_		90					95		See #10	
		11031	(XXX)) Asn	Lys((XXX)	Ala	Gln	Glu	Ile	Lys	Ser	Cys	Phe	$_{ m Lys}$	Leu	Gly	His	21.5	
		11032	•	_		100				7	105	_			-	110			on the	
		11034	Ile	Lys	Gly.	Lys	Lys	Gly	Ser	Glu	Arg	Arg	Val	Arg	Lys	Ile	Ser	Ser	1-000	
		11035	7		115			_	<u></u>	120	-	a 1	-		125	\	\ -1	7	Error	
	E>	11037	Gln		Thr	Lys	Asn	Leu	(XXX)	Arg	Arg	GIn	Pro	Pro	Asn (XXX) Ile	Arg	<u></u>	
		11038	<0105	130	D NO.	127			133					140					Summary	1
	11040 <210> SEQ ID NO: 417																			
	11041 <211> LENGTH: 74 11042 <212> TYPE: PRT																		Sheet.	
						homo	canio	ne'												
11043 <213> ORGANISM: homo sapiens 11045 <400> SEQUENCE: 417																				
		11047	Leu	-	Leu	Met	Gly	Arq	Leu	Ile	Tyr	Asn ((XXX)	Asn	Tyr	Leu	Phe	Tyr		
		11048	1		LICA	1100	5	111.9	LICU.	1.10	<i>x.j. x</i>	10			-1-	2000	15	-1-		
		11050	Lys	(XXX)	Asp	Ser	Ile	His	(XXX)	Gly	Arg		Leu	Glu	Val	Gln	Tyr	Thr		
		11051	17.0		E	20				V1	25	22.50				30	-1-			
		11053	Arg	XXX	Phe	Ile	Ser	Ser	Leu	His	Tyr	Asp	Cys	Glu	Phe	Pro	Tyr	Lys		
		11054	- · · · · · · · · · · · · · · · · · · ·		35					4.0	4		- 4		45			-		
	E>	11056	Leu (XXX	Thr	(XXX)	His	(XXX)	Lys	G1y	Asn(Xxx) Lys	11e	His	Phe	Tyr	Ile		
		11057		50		C.			55			- The second	-	60			-			
	E>	11059	His	Lys	Asn	Lys	Thr (Xxx	Pro ((xxx)	Glu	Thr								
				-				- Linesperson Contract		-										

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/673,840

DATE: 02/15/2001 TIME: 14:19:55

Input Set : A:\ES.txt
Output Set: N:\CRF3\02152001\1673840.raw

11060 65 70

```
<210> 391
                       Missing mandatory (220) to (223) features
<211> 3218
<212> DNA
                       to explain the "n's" at the circled locations in the sequence. See # 10 on
<213> homo sapiens
<400> 391
gcgaccacga gctggtgcat ccatcagtac ccttgccgga ctttccctta aagaaggaga 60
                                                                       the Error
ggatcagaaa gagataaaga ttgagccagc tcaggctgtg gatgaagtgg aacctctacc 120
tgaagactat tatacaagac cagtaaattt aacagaggta acaacccttc agcagcgtct 180
                                                                       Summary
gttacagect gaetteeage eagtetgtge tteacagete tateetegee acaaacatet 240
tctgatcaaa cggtccctgc gctgccgtaa atgtgaacat aatttgagca agccagaatt 300
taacccaacg tcaatcaaat tcaaaatcca gctggtcgct gtcaattata ttccagaagt 360
gagaatcatg tcaattccca accttcgcta catgaaggag agccaggtcc tcctgactct 420
tacaaatcca gttgagaacc tcacccatgt gactctcttc gagtgtgagg agggggaccc 480
tgatgatatc aacagcactg ctaaggtggt ggtgcctccc aaagagctcg ttttagctgg 540
caaggatgca gcagcagagt acgatgagtt ggcagaacct caagactttc aggacgatcc 600
tgacattata gccttcagaa aggccaacaa agtgggtatt ttcatcaaag ttacaccaca 660
gcgtgaggag ggtgaagtga ccgtgtgctt caagatgaag catgatttta aaaacctggc 720
agcccccatt cgccccattg aagaaagtga ccagggaaca gaagtcatct ggctcaccca 780
gcatgtggaa cttagcttgg gcccacttct tccttaaaag gttccactgg agggcagatc 840
ccaaaggaca gtatcaccgt aaacctgcgt taaaatgtgg aagctgctgc ttcattaggc 900
cttgtttata acgatgtacc catgcactac ggaattctat tgctaagaaa gtgggagcat 960
aggcaaggca ttgggaacac agggtagctg ctgttgctct tgctctcacc cctgttgaca1020
ccagtaagtc tgtgtctccc tcactgaacc ctgcacgttg agtaacagca gcataattcc1080
atcctaggaa aggggatggg tgttccttgg aatggcattg tatttaccac ctgagaaact1140
ctgtactgtc tcttgatctg atctcactaa ggatcacaat gtcacagatg aaacttaaat1200
gataacccaa aggtagacct gctgttaatg atccagcatt ggtcacaatg taccaactgc1260
tttctgcatt ccgttaaata tcatctaaca gtctaaaaca tatcccttca ttgccataat1320
ggctgccatt ttgccataga tttccatata actgaaaaac tgaattgtca ctttatcttt1380
agtatcatga tgattggaaa aacctgtgaa gttgttaagg cactctcatt tgccctcttt1440
ttctaagtga atacaggaca cgtattagtt gttcttaann nnnnnnnnn nnnnnnnnn 1500
```

XF.Y. 1.

<u>Please Note:</u>

Use of n and/or Xaa have been detected in the Sequence Listing. Please review the Sequence Listing to ensure that a corresponding explanation is presented in the <220> to <223> fields of each sequence which presents at least one n or Xaa.

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/673,840

DATE: 02/15/2001 TIME: 14:19:56

Input Set : A:\ES.txt

Output Set: N:\CRF3\02152001\1673840.raw

L:11 M:270 C: Current Application Number differs, Replaced Current Application Number L:12 M:271 C: Current Filing Date differs, Replaced Current Filing Date L:132 M:283 W: Missing Blank Line separator, <400> field identifier L:133 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (5) SEQUENCE: L:319 M:283 W: Missing Blank Line separator, <400> field identifier L:320 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (10) SEQUENCE: L:323 M:283 W: Missing Blank Line separator, <400> field identifier L:324 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (11) SEQUENCE: L:449 M:283 W: Missing Blank Line separator, <400> field identifier L:450 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (15) SEQUENCE: L:453 M:283 W: Missing Blank Line separator, <400> field identifier L:454 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (16) SEQUENCE: L:530 M:283 W: Missing Blank Line separator, <400> field identifier L:531 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (19) SEQUENCE: L:716 M:283 W: Missing Blank Line separator, <400> field identifier L:717 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (28) SEQUENCE: L:1039 M:283 W: Missing Blank Line separator, <400> field identifier L:1040 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (41) SEQUENCE: L:1043 M:283 W: Missing Blank Line separator, <400> field identifier L:1044 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (42) SEQUENCE: L:1083 M:283 W: Missing Blank Line separator, <400> field identifier L:1084 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (45) SEQUENCE: L:1140 M:283 W: Missing Blank Line separator, <400> field identifier L:1141 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (49) SEQUENCE: L:1444 M:283 W: Missing Blank Line separator, <400> field identifier L:1445 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (64) SEQUENCE: L:1470 M:283 W: Missing Blank Line separator, <400> field identifier L:1471 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (66) SEQUENCE: L:1486 M:283 W: Missing Blank Line separator, <400> field identifier L:1487 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (68) SEQUENCE: L:1533 M:283 W: Missing Blank Line separator, <400> field identifier L:1534 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (70) SEQUENCE: L:1537 M:283 W: Missing Blank Line separator, <400> field identifier I:1538 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (71) SEQUENCE: L:1611 M:283 W: Missing Blank Line separator, <400> field identifier L:1612 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (74) SEQUENCE: L:1636 M:283 W: Missing Blank Line separator, <400> field identifier L:1637 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (76) SEQUENCE: L:1741 M:283 W: Missing Blank Line separator, <400> field identifier L:1742 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (81) SEQUENCE: L:1785 M:283 W: Missing Blank Line separator, <400> field identifier L:1786 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (84) SEQUENCE: L:1815 M:283 W: Missing Blank Line separator, <400> field identifier L:1816 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (87) SEQUENCE: L:1842 M:283 W: Missing Blank Line separator, <400> field identifier L:1843 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (89) SEQUENCE: L:1869 M:283 W: Missing Blank Line separator, <400> field identifier L:1870 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (91) SEQUENCE:

VERIFICATION SUMMARY
PATENT APPLICATION: US/09/673,840

DATE: 02/15/2001 TIME: 14:19:56

Input Set : A:\ES.txt

Output Set: N:\CRF3\02152001\1673840.raw

L:3227 M:283 W: Missing Blank Line separator, <400> field identifier L:3228 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (130) SEQUENCE: L:3438 M:283 W: Missing Blank Line separator, <400> field identifier L:3439 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (140) SEQUENCE: L:3442 M:283 W: Missing Blank Line separator, <400> field identifier L:3443 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (141) SEQUENCE: L:3446 M:283 W: Missing Blank Line separator, <400> field identifier L:3447 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (142) SEQUENCE: L:4427 M:283 W: Missing Blank Line separator, <400> field identifier L:4428 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (175) SEQUENCE: $L:4431\ M:283\ W:$ Missing Blank Line separator, <400> field identifier L:4432 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (176) SEQUENCE: L:4435 M:283 W: Missing Blank Line separator, <400> field identifier L:4436 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (177) SEQUENCE: L:4439 M:283 W: Missing Blank Line separator, <400> field identifier L:4440 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (178) SEQUENCE: L:4443 M:283 W: Missing Blank Line separator, <400> field identifier L:4444 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (179) SEQUENCE: L:4447 M:283 W: Missing Blank Line separator, <400> field identifier L:4448 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (180) SEQUENCE: L:4565 M:283 W: Missing Blank Line separator, <400> field identifier L:4566 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (185) SEQUENCE: L:4569 M:283 W: Missing Blank Line separator, <400> field identifier L:4570 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (186) SEQUENCE: L:4573 M:283 W: Missing Blank Line separator, <400> field identifier L:4574 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (187) SEQUENCE: L:4777 M:283 W: Missing Blank Line separator, <400> field identifier L:4778 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (197) SEQUENCE: L:4781 M:283 W: Missing Blank Line separator, <400> field identifier L:4782 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (198) SEQUENCE: L:4785 M:283 W: Missing Blank Line separator, <400> field identifier L:4786 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (199) SEQUENCE: L:5607 M:283 W: Missing Blank Line separator, <400> field identifier L:5608 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (230) SEQUENCE: L:5611 M:283 W: Missing Blank Line separator, <400> field identifier L:5612 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (231) SEQUENCE: L:5615 M:283 W: Missing Blank Line separator, <400> field identifier L:5616 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (232) SEQUENCE: L:5665 M:283 W: Missing Blank Line separator, <400> field identifier L:5666 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (234) SEQUENCE: L:5669 M:283 W: Missing Blank Line separator, <400> field identifier L:5670 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (235) SEQUENCE: $L:5673\ M:283\ W:$ Missing Blank Line separator, <400> field identifier L:5674 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (236) SEQUENCE: L:5734 M:283 W: Missing Blank Line separator, <400> field identifier L:5735 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (240) SEQUENCE: L:5738 M:283 W: Missing Blank Line separator, <400> field identifier L:5739 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (241) SEQUENCE: L:5742 M:283 W: Missing Blank Line separator, <400> field identifier

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/673,840

DATE: 02/15/2001 TIME: 14:19:56

Input Set : A:\ES.txt

Output Set: N:\CRF3\02152001\I673840.raw

L:5743 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (242) SEQUENCE: L:5826 M:283 W: Missing Blank Line separator, <400> field identifier L:5827 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (245) SEQUENCE: L:5830 M:283 W: Missing Blank Line separator, <400> field identifier L:5831 M:300 W: (50) Intentionally skipped Sequence, : Sequence Id (246) SEQUENCE: L:9955 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:391 L:9955 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:391 L:9955 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:391 L:9955 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:391 L:9955 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:391 L:9956 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:391 L:9956 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:391 L:9956 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:391 L:9956 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:391 M:340 Repeated in SegNo=391 L:10173 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:396 L:10173 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:396 L:10173 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:396 L:10173 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:396 L:10173 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:396 L:10193 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10193 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10193 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10193 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10193 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:397 L:10195 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10195 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10195 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10195 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 M:340 Repeated in SegNo=397 L:10197 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10197 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10197 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10197 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10198 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10198 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10198 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10198 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10199 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10199 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10199 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10199 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10200 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10200 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10200 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10200 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10201 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10201 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10201 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397

VERIFICATION SUMMARY

PATENT APPLICATION: US/09/673,840

DATE: 02/15/2001 TIME: 14:19:56

Input Set : A:\ES.txt

Output Set: N:\CRF3\02152001\1673840.raw

 $\rm L\!:\!10201~M\!:\!258~W\!:$ Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10202 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10202 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10202 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10202 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10203 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10203 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:10203 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:397 L:10203 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:397 L:10204 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:397 L:10204 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:397 L:11013 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:3 L:11016 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:1 L:11019 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:3 L:11022 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:3 L:11025 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:2 L:11028 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:3 L:11031 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:2 L:11037 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:2 L:11047 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:1 L:11050 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:2 L:11053 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:1 L:11056 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:4 L:11059 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:2